**Title:** Tree height and hydraulic traits shape growth responses across droughts in a temperate broadleaf deciduous forest

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Text	word count	other	n
Total word count (excluding summary, references and legends)	currently ~6442 (strict limit 6,500)	No. of figures	2 (both colour)
Summary	currently 200 (limit 200)	No. of Tables	5
Introduction	currently $\sim 1339$	No of Supporting Information files	6
Materials and Methods	currently $\sim 1604$		
Results	currently $\sim 1558$		
Discussion	currently ~1866		
	(limit $30\%$ of		
	total (not strict),		
	or 1950 if		
	manuscript		
	reaches word		
	limit)		
Acknowledgements	currently $\sim 73$		

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## Summary

- As climate change is driving increased drought in many forested regions around the world, mechanistic understanding of factors conferring drought resistance in trees is increasingly important. However, it remains unclear how tree size and species' traits interactively shape tree growth responses during droughts.
- In this study, we analyze tree-ring records for twelve species of an oak-hickory forest representing 97% of woody productivity in the 25.6-ha ForestGEO plot in Virginia (USA) to determine how tree size, microhabitat, and species' traits interactively shape drought responses across the three strongest droughts over a 60-year period (1950 2009).
- Individual-level drought resistance decreased with tree height, whereas crown position and soil water availability explained little variation. Resistance was lower among species whose leaves had a higher percent loss of area upon dehydration and lost turgor at less negative water potential. However, there was substantial variation in the best predictor variables across the three drought periods. See comment 1 and 2
- We conclude that height—as opposed to canopy position, soil water availability, or species' traits—is the primary factor through which tree size mediates drought response, that leaf hydraulic traits are useful indicators of drought sensitivity, and that directions but not strengths of responses are consistent across droughts.

Key words: canopy position; drought; Forest Global Earth Observatory (ForestGEO); leaf hydraulic traits; temperate broadleaf deciduous forest; tree growth; tree height; tree-ring [5-8] Five to eight key words (in alphabetical order). Words that are in the title can, and should, be among these. See comment 3

### Introduction

See comment 4 Forests globally play a critical role in climate regulation (Bonan, 2008), yet there remains enormous uncertainty as to how the terrestrial carbon (C) sink, which is dominated by forests, will respond to climate change (Friedlingstein et al., 2006). An important aspect of this uncertainty lies with physiological responses of trees to drought (Kennedy et al., 2019). In many forested regions around the world, the risk of severe drought is increasing (Trenberth et al., 2014; Cook et al., 2014, 2015; Mankin et al., 2019), sometimes despite increasing precipitation (Intergovernmental Panel on Climate Change, 2015). Droughts resulting from global change have been affecting forests worldwide (Allen et al., 2010), and are expected to continue severely impacting forests (Anderegg et al., 2013; Adams et al., 2009; Bréda et al., 2006; Hartmann et al., 2018; Allen et al., 2015; Abrams and Nowacki, 2016; Adams et al., 2017). During drought, larger trees generally suffer greater reductions in growth and increased mortality rates (e.g., Bennett et al., 2015; Stovall et al., 2019), resulting in disproportionate impacts on forest C storage (Meaker et al., 2018). As a result, forest drought responses stand to strongly impact forest feedbacks to climate change (McDowell et al., 2016; Charney et al., 2016). Yet, accurate characterization of drought responses remains a modeling challenge (Kennedy et al., 2019)—in part because some of the underlying mechanisms remain unclear (REF). Understanding forest responses to drought requires increased functional understanding of how tree size, microhabitat, and species' traits jointly confer individual-level vulnerability or resistance, and the extent to which their influence is consistent across droughts.

One fundamental question regarding forest responses to drought is what drives the observed tendency for large trees to suffer more during drought. Bennett et al. (2015) was the first to show larger growth reductions for larger trees on a global scale, and numerous subsequent studies have reinforced this finding (e.g., Stovall et al. (2019); Hacket-Pain et al. (2016)]. However, they quantified tree size based on DBH, which has no direct mechanistic response. Their study proposed two major mechanisms-besides the tendency for bark beetles to preferentially attack larger trees (Pfeifer et al., 2011)-for the observed greater drought growth reductions of large trees see comment 5. First, Bennett et al. (2015) found taller trees have a greater biophysical challenge of lifting water greater distances against the effects of gravity and friction (McDowell et al., 2011; McDowell and Allen, 2015; Ryan et al., 2006; Couvreur et al., 2018), especially during drought (Zhang et al., 2009). Second, larger trees may have lower drought resistance because of their higher crown positions, where they are exposed to higher solar radiation, greater wind speeds, lower humidity, and lower CO 2 concentrations [Koike et al. (2001); Kunert et al. (2017) **REFS-KAT**). Subcanopy trees subsequently tend to fare better specifically due to the benefits of a buffered environment (Pretzsch et al., 2018). Potentially counteracting the biophysical challenges faced by large trees, their larger root systems confer an advantage in terms of allowing greater access to water; however, it appears that this effect is usually insufficient to offset the costs of height and/or crown exposure (Bennett et al., 2015). see comment 6 In addition to what was described by Bennett et al. (2015), a final mechanism that could mediate tree size-related responses to drought is how hydraulic traits are distributed with respect to size (Meakem et al., 2018; Liu et al., 2019). The pattern observed by Bennett et al. (2015) could be caused if the larger size classes were dominated by species less adapted to handle drought, be it through avoidance, resistance, or resilience see comment 7. Alternatively, larger size classes may be dominated by species that are inherently better adapted to greater biophysical challenges—as is the case in tropical moist forests of Panama, where larger size classes contain greater proportions of deciduous species (Condit et al., 2000; Meakem et al., 2018). Understanding the mechanisms driving more severe suffering of larger trees during drought will require sorting out the interactive effects of height, canopy position, soil water availability, and species' traits.

A second fundamental question regarding forest responses to drought is how species' traits – alone and in interaction with tree size – influence drought response (Elliott et al., 2015; Martin-Benito and Pederson, 2015). Separating species by ring porosity provides a first step toward identifying sensitivity, with diffuse-porous species tending to be more sensitive than ring-porous species (Kannenberg et al., 2019; Elliott et al., 2015; Friedrichs et al., 2009), but additional insight similar to Roskilly et al. (2019) is needed to better predict sensitivity at species or individual levels. Commonly-measured traits including wood density (WD) and leaf mass per area (LMA) have been linked to drought responses in temperate deciduous forests (Abrams, 1990; Guerfel et al., 2009; Hoffmann et al., 2011) and other forest biomes around the world (Greenwood et al., 2017). However, the direction of response is not always consistent; wood density correlated negatively with drought tree performance in a broadleaf deciduous forest in the southeastern United States (Hoffmann et al., 2011) but positively or with no effect at a global scale [Greenwood et al. (2017); Anderegg et al. (2018)). Hydraulic traits including water potentials at which percent loss conductivity surpass a certain threshold (P50, P80, P88) and hydraulic safety margin tend to be more successful at predicting drought performance (Anderegg et al., 2018) but are time-consuming to measure and therefore infeasible for predicting or modeling drought responses in highly diverse forests (e.q., in the tropics). More rapidly measurable leaf hydraulic traits with direct linkage to plant hydraulic function are emerging with potential to explain greater variation in plant distribution and function than the more commonly-measured WD and LMA (Medeiros et al., 2019). These include traits such as the fraction of conductivity lost to embolism, is another relevant metric (percent loss conductivity) [REF??]. leaf area shrinkage upon dessication ( $PLA_{dry}$ ; (Scoffoni et al., 2014)), and the leaf water potential at turgor loss point  $(\pi_{tlp})$ , i.e., the water potential at which leaf wilting occurs (Bartlett et al., 2016). The abilities of both  $PLA_{dry}$  and  $\pi_{tlp}$  to explain tree performance under drought remains untested.

A final central question regarding forest responses to drought is whether tree size and species' traits have similar influence across different drought events, or whether that influence is more strongly predicted by community-level responses to variable drought severity, duration, and timing based on tree size and traits. Tree growth responses vary with drought characteristics such as timing and atmospheric demand (D'Orangeville et al., 2018), but the question of how tree size and species' traits impact growth responses across droughts still remains. While tree-ring studies provide long-term records of tree responses to multiple droughts (e.g., (Lloret et al., 2011; D'Orangeville et al., 2018) **REFS**), they don't test for differential trait effects across periods of water shortage (D'Orangeville et al., 2018) and generally see comment 8 focus on species-level responses, and do not consider the roles of tree size and microenvironment. The ecological field-based studies that have shaped our understanding of the role of tree size and microenvironment in forest drought responses generally examine only a single drought and tend to focus disproportionately on extreme droughts with dramatic impacts (e.g., [Allen et al. (2010); Bennett et al. (2015); Stovall et al. (2019); Anderegg et al. (2016)). Thus, our knowledge of forest responses to more modest but frequent droughts - e.g., those with historical return intervals on the order of one to two decades - remains limited. While the tendency for larger trees to experience more severe drought certainly predominates the literature (Bennett et al., 2015), there are exceptions (e.g., Suarez et al., 2004; Zang et al., 2012; McDowell et al., 2008). There is also evidence that the degree to which larger trees are more impacted by drought increases with the severity of drought conditions (Bennett et al., 2015; Stovall et al., 2019). [Are there any studies showing interactions of drought type with traits?] see

**comment 9** Thus, while we expect many of the mechanisms shaping drought responses to be universal, we have little understanding of how tree size and traits interact with drought characteristics.

To yield functional understanding of how tree size, microenvironment, and species' traits collectively shape drought responses, we test a series of hypotheses and associated specific predictions (Table 1) based on the combination of tree-ring records from three droughts (1966, 1977, 1999), species functional and hydraulic trait measurements, and forest census data from a 25.6-ha ForestGEO plot in Virginia (USA). First, we focus on the role of tree size and its interaction with microenvironment. We confirm that, consistent with most forests globally, larger-diameter trees tend to have lower drought resistance in this forest, which is in an ecoregion see comment 10 represented by only one study in (Bennett et al., 2015) (H1.0). We then test hypotheses designed to disentangle the relative importance of tree height (H1.1), crown exposure (H1.2), and soil water availability, which should be greater for larger trees in dry but not in perpetually wet microsites (H1.3). see comments 11,12. Second, we focus on the role of species' functional and hydraulic traits and their interaction with tree height. We hypothesize that drought resistance will be negatively affected by wood density (following observed patterns by e.g. Hoffmann et al. (2011)), and positively affected by specific leaf area and xylem architecture (i.e., ring or diffuse/ semi-ring porous), but that hydraulic leaf traits including  $PLA_{dry}$  and  $\pi_{tlp}$  will prove better predictors (H2.1). We then test whether these traits correlate with tree height (H2.2), potentially driving the observed tendency for taller trees to suffer more during drought after accounting for trait effects (H2.3). Finally, we focused on variability among droughts, asking how community resistance varied across droughts (H3.1) and whether the factors confirming vulnerability or resistance varied across droughts (H3.2).

## Materials and Methods

### Study site

Research was conducted at the 25.6 ha ForestGEO (Forest Global Earth Observatory) study plot at the Smithsonian Conservation Biology Institute (SCBI) in Virginia, USA (38°53'36.6"N, 78°08'43.4"W) (Bourg et al., 2013; Anderson-Teixeira et al., 2015a). SCBI is located in the central Appalachian Mountains at the northern edge of Shenandoah National Park. Elevations range from 273-338m above sea level with a topographic relief of 65m (Bourg et al., 2013). Climate is humid temperate, with mean annual temperature of 12.7°C and precipitation of 1005 mm during our study period (1960-2009; source: CRU TS v.4.01; Harris et al. (2014)). Dominant tree taxa within this secondary forest include *Liriodendron tulipifera*, oaks (Quercus spp.), and hickories (Carya spp.). see comment 13

## Data collection and preparation

All analysis beyond basic data collection was performed using R version 3.5.3 (R Core Team, 2019).

Within or just outside the ForestGEO plot, we collected data on a suite of variables including tree size, microenvironment, and species traits (Table 2). Destructive sampling was only carried out within the plot. see comment 14 The SCBI ForestGEO plot was censused in 2008, 2013, and 2018 following standard ForestGEO protocols, whereby all free-standing woody stems  $\geq$  1cm diameter at breast height (DBH) were mapped, tagged, measured at DBH, and identified to species (Condit, 1998). From this census data, we used measurements of DBH from 2008 to calculate historical DBH and data for all stems  $\geq$  10cm to analyze functional trait composition relative to tree height (all analyses described below). Census data, which were last updated in 2019, are available through the ForestGEO data portal (www.forestgeo.si.edu).

We analyzed tree-ring data (cambial growth increment) see comment 15 from 571 trees representing the twelve species contributing most to woody aboveground net primary productivity ( $ANPP_{stem}$ ), which together comprised 97% of study plot  $ANPP_{stem}$  between 2008 and 2013 (Helcoski et al., 2019) (Figure S1). Cores were collected at breast heigh (1.3m) in 2010-2011 or 2016-2017 using a 5mm increment borer. In 2010-2011, cores were collected from randomly selected live trees of species with at least 30 individuals  $\geq$  10cm DBH (Bourg et al., 2013). In 2016-2017, cores were collected from all trees found dead in the annual mortality census (Gonzalez-Akre et al., 2016). Cores were sanded, measured, and cross-dated using standard procedures, as detailed in (Helcoski et al., 2019). The resulting chronologies have been published in association with Helcoski et al. (2019): (ITRDB; GitHub/Zenodo). Ryan, have you heard back from ITRDB? We should also cite GitHub/Zenodo here.

For each cored tree, we combined tree-ring records and allometric equations of bark thickness to retroactively calculate DBH for the years 1950-2009. Prior DBH was estimated using the following equation:

$$DBH_Y = DBH_{2008} - 2 * \left[ \sum_{year=Y}^{2008} (r_{ring,Y} : r_{ring,2008}) - r_{bark,Y} + r_{bark,2008} \right]$$

Here, Y denotes the year of interest,  $r_{ring}$  denotes ring width derived from cores, and  $r_{bark}$  denotes bark thickness. Bark thickness was estimated from species-specific allometries based on the bark thickness data from the site (Anderson-Teixeira et al., 2015b). Specifically, we used linear regression equations on log-transformed data to relate bark thickness to diameter inside bark from 2008 data (Table S1), which were then used to determine bark thickness in the retroactive calculation of DBH.

Height measurements (n=1518 trees) were taken by several researchers between 2012 to 2019, and are archived in a public GitHub repository. Measurement methods included manual (Stovall et al., 2018a, NEON), digital rangefinders (Anderson-Teixeira et al., 2015b), and automatic, ground-based LiDAR (Stovall et al., 2018b). Rangefinders used either the tangent method (Impulse 200LR, TruPulse 360R) or the sine method (Nikon ForestryPro) for calculating heights. Both methods are associated with some error (Larjavaara and Muller-Landau, 2013), but in this instance there was no clear advantage of one or the other. Species-specific height allometries were developed (Table S2). For species with insufficient height data to create reliable species-specific allometries, heights were calculated from an equation developed using all height measurements. see comments 16,17

Crown positions were recorded in the field during the growing season of 2018 following the crown position protocol of Jennings et al. (1999), whereby positions were ranked as dominant, codominant, intermediate, or suppressed. As it was unfeasible to retroactively estimate crown position, we assumed that 2018 crown position was reflective of each tree's position over the past 60 years. While some trees undoubtedly changed position, an analysis of crown position relative to height (Fig. 2d) and height variance since the beginning of the study period indicated that change was likely slow (Fig. S3). see comment 18,19 Specifically, average tree height growth was confined to ~0.82m from 1966 to 1977, ~1.45m from 1977 to 1999, and ~1.97m from 1999 to 2018.

Topographic wetness index (TWI) was calculated using the dynatopmodel package in R (Figure S1) (Metcalfe et al., 2018). Originally developed by Beven and Kirkby (1979), TWI was part of a hydrological run-off model and has since been used for a number of purposes in hydrology and ecology (Sørensen et al.,

2006). TWI calculation depends on an input of a digital elevation model (DEM), and from this yields a quantitative assessment defined by how "wet" an area is, based on areas where run-off is more likely. From our observations in the plot, the calculation of TWI performed comparatively better at categorizing wet areas than the calculation of a distance matrix from a stream shapefile.

Hydraulic traits were collected at SCBI (Table 3) in August 2018. We sampled small sun-exposed branches see comment 20 from three individuals of each species in and around the ForestGEO plot. Sampled branches were re-cut under water at least two nodes above the original cut and re-hydrated overnight in covered buckets (opaque plastic bags) before measurements were taken. Rehydrated leaves (n=3 per individual) were scanned, weighed, dried at 60° C for ≥ 48 hours, and then re-scanned and weighed. Leaf area was calculated from scanned images using the LeafArea R package (Katabuchi, 2019). LMA was calculated as the ratio of leaf dry mass to fresh area. PLA was calculated as the percent loss of area between fresh and dry leaves. WD was calculated for  $\sim 1$ cm diameter stem samples (bark and pith removed) as the ratio of dry weight to volume. see comment 21 We used the rapid determination method of Bartlett et al. (2012) to estimate water potential at turgor loss point  $(\pi_{tlp})$ . Briefly, two 4mm diameter leaf discs were cut from each leaf, tightly wrapped in foil, submerged in liquid nitrogen, perforated 10-15 times with a dissection needle, and then measured using a vapour pressure osmometer (VAPRO 5520, Wescor, Logan, UT, USA). Osmotic potential  $(\pi_{osm})$  given by the osmometer was used to estimate  $(\pi_{tlp})$  using the equation  $\pi_{tlp} = 0.832\pi_{osm}^{-0.631}$  (Bartlett et al., 2012). Data and R scripts for hydraulic traits are available at [create new public GitHub repo for hydraulic traits, archive in Zenodo, give DOI]. see comment 22

To characterize how environmental conditions vary with height, data were obtained from the National Ecological Observation Network (NEON) tower located <1km from the study area. We used data on wind speed, relative humidity, and air temperature, all measured over a vertical profile **see comment 23**, for the years 2016-2018 (?). After filtering for missing and outlier values, the data was consolidated to represent the mean values per sensor height per day. The range of these means were then aggregated at a month scale.

## Identification of drought years

We identified droughts within the time period 1950-2009, defining drought (Slette et al., 2019) as events with widespread reductions in tree growth and anomalously dry peak growing season climatic conditions. Specifically, candidate drought years were defined as those where >25% of the cored trees experienced >30% reduction in basal area increment (BAI) relative to the previous 5 years, following the drought resistance (R) metric of (Lloret et al., 2011), and identified using the pointRes package (van der Maaten-Theunissen and van der Maaten, 2016) in R. see comment 24 Separately, we identified the years with driest conditions during May-August, which stood out in the analysis of (Helcoski et al., 2019) as the current-year months to which annual growth was most sensitive for trees at this site. We considered two metrics of moisture deficit: NOAA Divisional Data's Palmer Drought Severity Index (PDSI) and the difference between monthly potential evapotranspiration (PET) and precipitation (PRE). These data were obtained from the ForestGEO Climate Data Portal (https://github.com/forestgeo/Climate) in August 2018, with monthly PET and PRE sourced from Climatic Research Unit high-resolution gridded dataset (CRU TS v.4.01; Harris et al. (2014)). The driest years were identified through ranking mean May-August PDSI or [PET-PRE] for the time period from driest to wettest. see comment 25

Statistical Analysis

For each drought period, we calculated drought resistance (R) as the ratio of BAI during drought to the mean BAI over the five years preceding the drought (Lloret et al., 2011). Analyses focused on testing the predictions presented in Table 1, with either R or H as the response variable. see comment 26 The general statistical model for hypothesis testing was a mixed effects model (lme4 package from Bates et al. (2019)) with R as the response variable, tree nested within species as a random effect, and one or more independent variables as fixed effects. We used AICc (AICcmodavg package from Mazerolle and portions of code contributed by Dan Linden. (2019)) to assess model selection, and conditional/marginal R-squared to assess model fit.

Models were run for all drought years combined (with year as a fixed effect) and for each drought year independently. In order to determine the relative importance of the traits alone (H2.1), we first tested the predictor variables independently against both height and R given height's substantial influence. Variables were considered useful for predicting R when AICc was reduced by  $\geq 2$  relative to the appropriate null model lacking that variable (Table 4).

Once these beset predictor variables were determined, we calculated the best overall models for predicting R across all traits, each individual drought year, and all years combined. We compared models with all possible combinations of candidate variables and identified the full set of models within dAICc=2 of the top model (that with lowest AICc). These were counted as the "top multivariate models". When a variable appeared in all of these top models and the sign of the coefficient was consistent across models, this was counted as support for/ rejection of the associated prediction by the multivariate models. If the variable appeared in only some of the models, we considered this partial support. see comment 27

#### Results

### Focal droughts and their characteristics

In the 60-year period between 1950 and 2009, there were three droughts that met our criteria of anomalously dry climatic conditions coupled with substantial reductions in tree growth for at least some portions of the community: 1966, 1977, and 1999 (Figs. 1, S2). We excluded one year (1991) meeting the growth reduction criteria (26.5% of trees experienced >30% growth reduction, mean resistance= -13.8%) because this year was not among the driest in terms of May-August [PET-PRE] or PDSI (Table S3). Rather, the severity of growth reduction may be explained by defoliation by gypsy moths (*Lymantria dispar L.*) from approximately 1988-1995, which most stronly impacted *Quercus* spp. (*Cite Shenandoah paper, if accepted*). Climatically, these droughts included three of the five years between 1950 and 2009 with greatest moisture deficit (PET-PRE) during the peak growing season months of May-August, which are the months to which annual tree growth at this site is most sensitive (Helcoski et al., 2019). Specifically, 1966, 1977, and 1999 had mean MJJA PET-PRE of 83.37, 86.97, and 80 mm mo-1, respectively. The years 1964 and 2007 also ranked in the top five driest (PET-PRE =83.87 and 82.13 mm mo-1), but were not among the lowest in terms of PDSI and were not identified as a pointer year (Table S3)

The droughts differed in intensity and prior onset (Table S3, Fig. S2). In terms of intensity during the peak growing season of the focal year, the 1999 drought was the most intense (lowest PDSI) during May-July. It was surpassed in intensity in August by the 1966 drought, which was otherwise the second most intense during the peak growing season. The 1977 was the least intense throughout the growing season. All droughts increased in intensity from May through August, but differed in the months previous.

The 1966 drought was preceded by two years of moderate drought during the growing season and severe to extreme drought starting the previous fall. Similarly, the 1999 drought was preceded by severe to extreme drought starting the previous fall, but before that conditions were wetter than average until the previous June. The 1977 drought was preceded by 2.5 years of near-normal conditions, and was therefore the mildest of the three droughts by any measure.

Community-level tree growth responses to these droughts were modest, with mean resistance values of 0.86, 0.84, and 0.86 for 1966, 1977, and 1999, respectively (Fig. 1b). In each drought, roughly 30% of the cored trees suffered  $\geq$  30% growth reductions ( $R \leq$  0.7): 29.43% in 1966, 31.86% in 1977, and 26.81% in 1999. Some trees exhibited increased growth: (R > 1.0): 26.31% in 1966, 21.92% in 1977, and 25.57% in 1999.

### Tree size and drought resistance

Overall, our analysis confirmed the tendency for larger-diameter trees to show greater reductions in growth during drought (Bennett et al., 2015) (H1.0), although there was no significant effect for 1977 or 1999 individually (Tables 1, 4). The same held true for ln[H] as a univariate predictor (H1.1; Tables 1, 4). When combined with other predictor variariables in our multivariate models, the top models usually included an effect of ln[H], and its coefficient was consistently negative, as predicted (Tables 1, 5). We note that a non-significant positive correlation between ln[H] and R for 1999 became negative in the context of the multivariate models, again supporting H1.1 (Table 1).

Crown position was generally correlated with H, but with substantial variation (Fig. 2d). Crown position was a much poorer predictor of R than was H (Table 4), lending little overall support to H1.2 (Table 1). When considered alone, CP had a significant influence only in the 1966 drought, during which trees with dominant CP had the lowest R. When considered in conjunction with H, CP came out as a significant predictor only for the 1999 drought, during which suppressed and then intermediate trees had the lowest R. Crown position was included in almost half of the top models, with mixed results as to how R varied with CP (Table 5). Most commonly in these multivariate models, as in the univariate models (Table 4), the resistance of dominant trees was less than that of co-dominant trees but higher that of suppressed trees. Thus, CP was sometimes a useful predictor of R, but overall had a weak effect relative to that of H.

In the non-drought years for which we have vertical profiles in climate data (2016-2018), taller trees—or those in dominant crown positions— were generally exposed to higher evaporative demand during the peak growing season months (May-August; Fig. 2). Specifically, maximum daily wind speeds were significantly higher above the top of the canopy (40-50m) than within and below (10-30m) (Fig. 2a). Relative humidity was also somewhat lower during June-August, ranging from ~50-80 above the canopy and ~60-90% in the understory (Fig. 2b). Air temperature did not vary across the vertical profile (Fig. 2c).

Resistance was negatively correlated with ln[TWI] (Tables 4-5), negating the idea that trees in moist microsites would suffer less during drought. Nevertheless, we tested for a negative ln[H]\*ln[TWI] interaction (H1.3), which could indicate that smaller trees (with smaller rooting volume) have a greater tendency to suffer more in drier microenvironments with greater depth to the water table. H1.3 was rejected; the ln[H]\*ln[TWI] interaction was never signficant and had a consistently positive coefficient (Table 4).

Species' traits, height, and drought resistance

We partially support H2.1: Species' hydraulic traits -XP,  $PLA_{dry}$ , and  $\pi_{tlp}$ —were sometimes useful in explaining variation in drought responses, whereas LMA and WD were not (Tables 1,4,5). Specifically,

LMA and WD never significantly associated with R in the univariate models (all dAIC  $\leq$  0.22; Table 4), and therefore these were excluded as candidate variables for the full multivariate models. In contrast, XP,  $PLA_{dry}$ , and  $\pi_{tlp}$  all explained at least modest amounts of variation (dAIC > 1.0) in at least one drought (Table 4). Of these,  $PLA_{dry}$  was a strong predictor for the 1966 drought and overall, with consistently negative coefficients across all droughts.  $\pi_{tlp}$  never came out as significant (dAIC  $\leq$  2) in the univariate models but had a consistently negative coefficient (Table 4). Whereas ring-porous species had highest R overall and in the 1966 and 1999 droughts, they had lower R in 1977. Results were similar in the context of multivariate models (Table 5), except that  $PLA_{dry}$  had a small positive coefficient in the one 1999 model in which it was included.  $PLA_{dry}$ , and  $\pi_{tlp}$  were each included in the top overall model and in the top model for two of the three individual droughts (Table 5).

We reject H2.2, finding no evidence that taller trees tend to have traits associated with greater drought resistance. In part because of the large sample size ( $\mathbf{n}=\#$  trees–all individuals of our 12 focal species  $\geq 10$  cm in 25.6 ha), there were very significant (p<0.0001) correlations of H with all species' traits (see Table S4). However, the correlation only matched the predicted direction (*i.e.*, more drought-resistant traits associated to taller trees) in the cases of WD and LMA, which were not useful predictors of R. Furthermore, although correlations were statistically significant, trait variation within each height class overwhelmed any vertical trends (Fig. 2e-f).

We support the hypothesis (H2.3) that the observed tendency for larger trees to have greater growth reductions during drought (lower R) is driven by height itself, as opposed to more drought-sensitive traits in larger trees (Tables 1,5). As discussed above, there was little meaningful variation in traits with height at the community level. When ln[H] and hydraulic traits were considered together in multivariate models, the effect of ln[H] on R was consistently negative (Table 5)—reversing a non-signficant positive ln[H]-R correlation in the univariate model for the 1999 drought (Table 4).

## Responses across droughts

We reject the hypothesis (H3.1) that overall community responses varied across droughts. Within the context of mixed effects models, there were no significant differences in R across drought years (Table 4). This is consistent with the observation that the distribution of R values was similar across droughts (Fig. 1b).

We mostly reject the hyopthesis (H3.2) that directions of responses varied across droughts. In the majority of cases, response directions were consistent across droughts in both univariate and multivariate models (Tables 1,4,5). However, there were a few exceptions—most commonly in the categorical variables (CP and XP) but also for  $PLA_{dry}$  in one multivariate model for the 1999 drought (Tables 4, 5). These differences may very well be random, as opposed to statistically meaningful. Among the univariate models, there was no instance where predictor variables significantly improved the models of two different droughts ( $dAIC \ge 2$ ), but with contrasting coefficients (Table 4). Among the multivariate models, CP was not consistently in the top models for any drought (Table 5), and  $PLA_{dry}$  only appeared with a positive coefficient in the weakest of six models for the 1999 drought (contrasting with a negative coefficient in the univariate model; Table 4). The difference most likely to be real is that ring-porous species had lower R than diffuse- and semi-ring- porous species in the 1977 drought, contrasting with higher resistance in 1966 and 1999 (Tables 4,5), but note that XP was not a significant predictor on its own for the 1977 drought.

We support the hypothesis (H3.3) that the strength of predictor variables was different across the droughts

(Tables 1,4,5). For instance, ln[H] and  $PLA_{dry}$  had much stronger negative effects in 1966 than in the other two years, ln[TWI] had the strongest negative effect in 1977, and XP (higher R among ring-porous trees) was strongest in 1999 (Tables 4,5).

## Discussion

Our results reveal how tree size, microhabitat, and hydraulic traits shaped tree growth responses across three droughts in a temperate deciduous forest (Table 1). The tendencey for larger trees to suffer more, observed here as in forests around the world (Bennett et al., 2015), was driven primarily by their height. There was a marginal additional effect of crown exposure, with the most exposed and the most suppressed trees suffering most-consistent with observations of both greater drought sensitivity of exposed trees (e.g., (Suarez et al., 2004); (Scharnweber et al., 2019)) and greater sensitivity of suppressed and crowded individuals (**REFS**). There was no evidence that soil water availability increased drougth resistance; in contrast, trees in wetter topographic positions suffered more (consistent with (Zuleta et al., 2017)), and larger rooting volume provided no advantage in the drier microenvironments. The lower drought resistance of larger trees was not driven by any tendency for the canopy to be dominated by more drought-sensitive species. Drought sensitivity was not consistently linked to species' LMA, WD, or xylem architecture. However, drought resistance was negatively correlated with the leaf hydraulic traits  $(PLA_{dru}, \pi_{tlp})$  in the top overall model and the top models for two of the three individual droughts, which is physiologically logical ((Scoffoni et al., 2014); (Bartlett et al., 2016); (Medeiros et al., 2019)) but scientifically novel in that  $PLA_{dry}$  and  $\pi_{tlp}$  have not previously been linked to drought growth responses. The direction of responses was mostly consistent across droughts, indicating that they were driven by fundamental physiological mechanisms; however, the strengths of each predictor varied across droughts, indicating that specific drought characteristics interact with tree size, microenvironment, and traits to shape which individuals suffer most. These findings significantly advance our knowledge of the factors that confer vulnerability or resistance on trees during drought.

The droughts considered here were of similar severity (Fig. 1b) and fairly moderate; droughts of this magnitude have occurred with an average frequency of approximately one per 10-15 years (Fig. 1a, Helcoski et al. (2019)). Therefore, we except that most species are adapted, and individual trees acclimatized, to survive droughts of this nature. While the majority of trees experienced reduced growth, a substantial portion had increased growth (Fig. 1b), underlining the fact that these droughts did not induce extreme stress on the entire forest. It is likely for this reason, combined with the fact that many factors other than climate affect tree growth in closed-canopy forests, that our best models characterize only a modest amount of variation: 11-13% for all droughts combined, and 21-26% for each individual drought (Table 5). Methodologically, the moderate nature of these droughts is an advantage because our analysis considers only trees that survived all of these droughts, and we lack information on the trees that were killed. These are likely to be relatively modest in number, and local forest monitoring data stretching back to the late 1980s confirms that the 1999 drought did not trigger major declines in tree abundance or biomass (Anderson-Teixeira et al., in revision). Thus, the droughts considered here are substantially weaker than those that have triggered massive tree die-off (e.g., (Allen et al., 2010)), many of which have shaped our understanding about the role of tree size (Bennett et al., 2015; Stovall et al., 2019) and to some extent-traits underlying drought response mechanisms (Greenwood et al. (2017); Anderegg et al. (2016)). Nevertheless, our results are consistent with findings from more extreme droughts.

Our analysis indicates that height-as opposed to canopy position or soil water availability-is the primary factor through which tree size mediates drought response. Taller trees face inherent biophysical challenges in lifting water a greater distance against the effects of gravity and friction (Ryan et al., 2006; McDowell and Allen, 2015; McDowell et al., 2011; Couvreur et al., 2018). Vertical gradients in stem and leaf traits-including smaller and thicker (higher LMA) leaves, more negative P50, and lower hydraulic conductivity at greater heights (Couvreur et al., 2018; Koike et al., 2001; McDowell et al., 2011)-make it biophysically possible for trees to become tall (Couvreur et al., 2018), yet height becomes a liability when drought incurs additional hydraulic challenges. Taller trees also face different microenvironments (Fig. 2a-b), in part because they are more likely to be in dominant canopy positions (Fig. 2d). Even under non-drought conditions, evaporative demand increases with tree height in taller trees are more closely coupled to the atmosphere (REFS- Jarvis 1984?; (Bretfeld et al., 2018)). Exposed canopy leaves reach higher temperatures (Smith & Nobel 1977), particularly during drought when solar radiation tends to be higher and less water is available for evaporative cooling of the leaves. Furthermore, daytime CO 2 concentrations tend to decrease with height (Koike et al., 2001), implying that water costs of  $CO_2$  uptake increase with height. Correlation between height and canopy position (Fig. 2d) makes it challenging to disentangle the relative importance of height per se from microenvironment. However, signficant decoupling between height and canopy position can be introduced by the configuration of neighboring trees (Fig. 2d) (Muller-Landau et al., 2006), and we show that height is a far stronger predictor of drought response than crown position (Tables 1,4,5). Our analysis does have the limitation that canopy positions were recored in 2018 and undoubtedly changed for some trees since the 1960s, and we note that CPbecame an increasingly poor predictor moving from 1999 back to 1966 (Table 4). However, because trees would generally advance towards more dominant positions as they grow and as neighbors die, changing canopy positions would bias against the acceptance of H1.2. The implication is that dominant crown positions did have a marginally negative influence on R, which makes sense in light of the vertical environmental gradients described above and agrees with previous studies showing greater drought sensitivity in more exposed trees ((Suarez et al., 2004); (Scharnweber et al., 2019)). It is safe to assume that currently suppressed trees have always been suppressed, and their relatively low R (after correcting for height effects) is real, which is consistent with analyses showing that suppressed—and particularly crowded—trees can suffer disproportionately during drought (REFS- Alan?). The observed height-sensitivity of R, together with the apparent lack of importance of soil water availability (H1.3), agrees with the concept that physiological limitations to transpiration under drought shift from soil water availability to the plant-atmosphere intreface as forests age (Bretfeld et al., 2018), such that tall trees-particularly the very tallest-are the most sensitive in mature forests. Additional research comparing drought responses of young and old forest stands, along with short and tall isolated trees, would be valuable for more clearly disentanging the roles of tree height and crown exposure.

The development of tree-ring chronologies for all dominant tree species at our site (Helcoski et al., 2019) made it possible to compare historical drought responses across 12 species and their associated traits at a single site for the first time (**verify- Neil, Alan**). Concerted measurement of leaf hydraulic traits of emerging importance (Scoffoni et al., 2014; Bartlett et al., 2016; Medeiros et al., 2019) allowed novel insights into the role of hydraulic traits in shaping drought response. The finding that  $PLA_{dry}$  and  $\pi_{tlp}$  can be useful for predicting drought responses—with both appearing in the top overall model and two of the three top models for each drought (Table 5)—is consistent with studies demonstrating that these are physiologically meaningful traits linked to species distribution along moisture gradients (Medeiros et al.,

2019) (MORE REFS–KAT/NOBBY/LAWREN). It is scientifically exciting in that this finding indicates that  $PLA_{dry}$  and  $\pi_{tlp}$ , which can be measured relatively easily (Bartlett et al., 2012; Scoffoni et al., 2014), hold promise for predicting drought growth responses across species. The importance of linking species' traits to drought responses increases with tree species diversity; whereas it is feasible to study drought responses for all dominant species in most boreal and temperate forests (e.g., this study), this becomes difficult to impossible for diverse tropical forests, where linking hydraulic traits to drought responses would be invaluable for forecasting how little-known species and whole forests will respond to future droughts (REFS?–KAT/NOBBY/LAWREN).

Our analysis of hydraulic traits focused on species-level comparisons and did not characterize the role of variation with height. As noted above, leaves found higher on a tree tend to have more hydraulically conservative traits, and therefore we would expect that average leaf characteristics of an individual tree would scale with its crown height, with taller individuals having on average more drought-resistant traits. If vertical trends for  $PLA_{dry}$  and  $\pi_{tlp}$ , which have not been characterized (LAWREN, IS THIS TRUE?), follow the general pattern of increasing leaf drought resistance with height, our analysis would be biased in a conservative direction when assessing  $H2.3^*$  (Table 1)–i.e., accounting traits on an individual level should result in a stronger negative effect of ln[H]. Further characterization of leaf hydraulic traits in relation to height and crown exposure would be valuable for enhancing our understanding of the interactive effects of tree height and traits on drought responses.

Although the physiological mechanisms discussed above lead to generally consistent directions of growth responses to tree height and hydaulic traits across droughts, indicating the universality of the underlying mechanisms, the relative importantance of the drivers varied widely across droughts, indicating an interaction with drought characteristics (Tables 4-5). Although there were not significant differences in Ramong the drought years, R tended to be somewhat lower in 1966 (Fig. 1, Table 4). Height and dominant canopy position had the stronger negative effects in this drought than in the others (Tables 4-5), which could potentially be linked to the fact that this was the longest-lasting drought and and that height becomes a stronger predictor of mortality as the severity of the drought increases (Stovall et al., 2019). In 1977, which was the weakest drought in terms of PDSI (Fig. S2, Table S3), ring-porous species had lower R, contrasting with their higher R in the other two droughts and perhaps indicating that ring-porosity provides less of an advantage in terms of drought adaptation under less extreme droughts, consistent with Elliott et al. (2015). In the 1999 drought, which was the most intense in terms of growing season PDSI and the second-longest in duration, xylem architecture was the most useful predictor, with diffuse-porous species suffering most. Thus, responses differed with the nature of each drought. Of course, site characteristics also define the nature of droughts, and comparisons of size and trait effects across sites—and across more droughts at a single site-would be of great value to elucidating the mechanisms through which drought characteristics interact with driver variables to shape tree growth responses.

As climate change drives increasing drought in many of the world's forests (Trenberth et al., 2014; Intergovernmental Panel on Climate Change, 2015), the fate of forests and their climate feedbacks will be shaped by the biophysical and physiological drivers observed here. Large trees have been suffering disrpoportionately in forests around the world (Bennett et al., 2015; Stovall et al., 2019), and we here show that this is primarily driven by their height, with some contributions from canopy position. The distinction is important because it suggests that height *per se* makes trees vulnerable, even if their crowns are somewhat protected by neighbors, whereas solitary trees or the dominant trees in young regrowth forests

should be less vulnerable. Considering just height and crown position, this would suggest that mature forests would be more vulnerable to drought than young forests with short trees; however, soil water availability may limit the young forests (Bretfeld et al., 2018), and species traits often shift as forests age. Pioneer species at our site (*Liriodendron tulipifera*, *Quercus spp.*, *Fraxinus americana*) have a mix of traits conferring drought tolerance and resistance (Table 3), and further research on how hydraulic traits and drought vulnerability change over the course of succession would be valuable for getting at the very significant question of whether and how drought tolerance changes as forests age. In the meantime, the results of this study advance our knowledge of the factors conferring drought vulnerability and resistance in a mature forest, opening the door for more accurate forecasting of forest responses to future drought.

## Acknowledgements

Thanks to all researchers who helped to collect the data used here, in particular Jennifer C. McGarvey, Jonathan R. Thomspon, and Victoria Meakem for original collection and processing of cores. Thanks also to Camila Medeiros for guidance on hydraulic and functional trait measurements and to Edward Brzostek's lab for collaboration on leaf sampling. \*\* others??\*\* This study was funded by the Smithsonian's Forest Global Earth Observatory (ForestGEO), a Virginia Native Plant Society grant to KAT and AJT, and support from the Harvard Forest and National Science Foundation which supports the PalEON project (NSF EF-1241930) for NP. (Neil, this is copied from Ryan's paper. accurate here? coauthors, other funding?)

### **Author Contribution**

KAT, IM, and AT designed the research. Tree-ring chronologies were developed by RH under guidance of AT and NP. Trait data was collected by IM, JZ under guidance of NK and LS. Other plot data were collected by IM, AS, EGA, and NB under guidance of EGA and WM. Data analyses were performed by IM under guidance of KAT and VH. KAT and IM interpreted the results. IM and KAT wrote the first draft of manuscript, and all authors contributed to revisions. coauthors, please check/edit

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